

Europäisches Patentamt

European Patent Office

Office européen des brevets



D EP 1 174 941 A2

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication: 23.01.2002 Bulletin 2002/04 (51) Int Cl.7: H01M 10/40, H01G 9/02

- (21) Application number: 01117201.2
- (22) Date of filing: 16.07.2001
- (84) Designated Contracting States:
 AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU
 MC NL PT SE TR
 Designated Extension States:
 AL LT LV MK RO SI
- (30) Priority: 17.07.2000 JP 2000215521 26.09.2000 JP 2000291715
- (71) Applicant: MATSUSHITA ELECTRIC INDUSTRIAL CO., LTD.
 Kadoma-shi, Osaka 571-8501 (JP)

- (72) Inventors:
- Sonoda, Kumiko Hirakata-shi, Osaka 573-0036 (JP)
 - Ueda, Atsushi
 Osaka-shi, Osaka 546-0003 (JP)
- Iwamoto, Kazuya
 Sakai-shi, Osaka 593-8304 (JP)

80336 München (DE)

- (74) Representative: Leson, Thomas Johannes Alois, Dipi.-Ing. Tiedtke-Bühling-Kinne & Partner GbR, TBK-Patent. Bayarlaring 4
- (54) Non-aqueous electrolyte and electrochemical device comprising the same
- (57) A non-aqueous electrolyte is disclosed, which comprises a non-aqueous solvent and a solute represented by the general formula(1) MBRI PRPSH, wherein M is an alkeli metal atom or an ammonium group and R1 to R4 are each independently electron withdrawing groups or electron withdrawing atoms bound to B where at least one of R1 to R4 is other than a fluorine atom. The solute has a thermal stability substantially equal to

that of LIBF₄ and an anion portion having a high electronegativity, and easily dissociation into ion. Therefore, a non-aqueous electrolyte containing this solute has a high indisconductivity and is difficult to cause a great ation of a gas or deterioration in characteristics due to the decomposition of the solute, which occurs the decomposition of the solute, which occurs due to the decomposition of the solute.

Description

BACKGROUND OF THE INVENTION

- [0001] Electrochemical devices comprising a non-aqueous electrolyte have been used in a wide range of electric and electrolic appliances. Such electrochemical deviouse include, for example, non-aqueous electrolyte batteries such as a lithium battery, and capacitors such as an electric double layer capacitor and an electrolytic capacitor.
 - [0002] It is desirable that a non-aqueous electrolyte comprising a non-aqueous solvent and a solute has a high ionic conductivity. In order to obtain such a non-aqueous selvent new and a low viscosity is required. However, a non-aqueous solvent having a high dielectric constant usually has a strong polarity and therefore has a high viscosity. Thus, a mixed non-aqueous deterotyte containing a non-aqueous solvent with a high delectric constant such as ethylenc extraorbate (delectric constant solvent with a high delectric constant such as ethylenc extraorbate (delectric constant; 3.1) or ethyl methyl carbonate (delectric constant; 3.9) is commonly used.
- 15 [0003] As the solute constituting the non-equeous electrolyte, for example, LiPF₈, LiClO₄, LiBF₄, LiAlCl₄, LiSbF₆, LiSCN, LiCl, LiAF₆, LiCF₉SO₂, NLI or the like is used. Among them, LiPF₆ is most frequently used because it has a wide potential window and thus provides a non-equeous electroly he awing a light lonic conductivity.
 - [9004] While a non-aqueous electrolyte containing LIPE, has a high ionic conductivity of approximately 8.5 mS/cm at room temperature. I has a low themal stability Moreover, LIPE; is problemate in that it reacts sharply with moisture to decompose. HF, PF, or the like produced by the decomposition of LIPE, has an adverse effect on an electrochemical device. Accordingly, when an electrochemical device, which compresses a non-aqueous electrolyte containing LIPE, is used for an appliance generating a great amount of heat, the performance of the electrochemical device may deteriorate or a gas may be generated in setting the device.
 - [0005] Although the uses of LISO_CFS_ (CF_SO_g)/NL and the like have also been investigated, they are not proceeding towards practical utilization. This is because many of solutes containing an organic anion tend to cause a corrosion of a current collector made of aluminum, which is used in the positive electrode of a non-aqueous electrolyte battary.
 - [0006] On the other hand, LIBF_A, which is commonly used in lithium primary batteries, has a higher thermal stability as compared with LIPF_a and a potential window substantially equal to that of LIPF_A. However, a non-equeous electryble obtained by dissolving LIBF_A in a non-equeous electryble obtained by dissolving LIBF_A in a non-equeous selvent has a low ionic conductivity of about 2.9 mS/cm at room temperature. For this reason, LIBF_A is not used in likitim secondary batteries.
 - [9007] The reason why LIBF_a has a lower lonic conductivity as compared with LIFF_a is that only four fluorine atoms awing an electron withdrawing property are bound to one born atom in LIBF_a, whereas ski fluorine atoms are bounded to one phosphorus atom in LIFF_a. Since electronegativity depends on the number of the bound fluorine atoms, the electronegativity of the BF_a and role is lower than that of the FF_a anion. Therefore, it is concidered that LIEF_a is difficult to dissociate into a Li cation and a BF_a anion. Moreover, because the BF_a anion has a small lonic diameter, it tends to cause an association thereby to destrorizate the ionic conductivity of the electrowise.
 - [9008] The delectric loss in an electrolytic capacitor is greatly affected by the ionic conductivity of the non-aqueous electrolyte interposed between a negative electrode foil and a delectrol layer which comprises an oxide of aluminum or tantalum disposed on a positive electrode foil. When the dislectric loss is great, the frequency characteristics and the charge/dislectarge characteristics of the electrolytic capacitor are deteriorated. Nevertheless, a non-aqueous electrolyte accommodated in an electrolytic capacitor contains a solute which dissociates to produce a BF₄ anion, resulting in a creat diffectivit loss in the capacitor.

45 BRIEF SUMMARY OF THE INVENTION

- [0009] In view of the above situation, it is an object of the present invention to provide a non-aqueous electrolyte having a high thermal stability as well as a high ionic conductivity.
- [0010] More specifically, the present invention relates to a non-aqueous electrolyte comprising a non-aqueous solvent and a solute represented by the general formula (1): MBPIPRPRPR, wherein M is an alkali metal atom or an ammonium group and R¹ to R⁴ are each independently electron withdrawing groups or electron withdrawing atoms bound to B where at least one of R¹ to R¹ so either than a fluoring atom.
- [0011] In the general formula (1), it is preferred that at least one of R¹ to R⁴ is represented by the general formula (C). C_nF_{2m+1}. Where n is an integer of 1 to 4; or the general formula (3): C_mF_{2m+1}SO₂, where m is an integer of 1 to 4.
 Further, the reminders of R¹ to R⁴ are preferably fluorine atoms.
 - [0012] For the solute represented by the general formula (1), it is preferred to use at least one selected from the group consisting of LIB(CF₃)₄. LIBF(CF₃)₃. LIBF₃(CF₃)₂. LIBF₃(CF₃)₃. LIBF₃(CF₃SO₃)₄. LIBF(CF₃SO₃)₅. LIBF₃(CF₃SO₃)₅. LIBF₃(CF₃SO₃)

 $(C_9F_5SO_9)_2$ and LiBF₃ $(C_9F_5SO_9)$.

[0013] In the case where M is an ammonium group in the general formula (1), it is preferred that the ammonium group is represented by the general formula (4). PRPRPRPRP where P to PR are bound to M and are each independently nydrogen atoms, alkyl groups, alkenyl groups or any groups. In this case, the above solute is preferably at least one selected from the group constituting of (CH₃)₃MRGCF₃, (CH₃)MRFICF₃, (CH₃)₄MRFICF₃, (CH₃)MRFICF₃, (C

[0014] The present invention further relates to an electrochemical device comprising the above non-aqueous electrolyte.

[0015] Examples of the electrochemical device include a non-aqueous electrolyte battery, an electric double layer capacitor, an electrolytic capacitor and the like.

[0016] The non-aqueous electrolyte battery preferably comprises a positive electrode; a negative electrode; a separator interposed between the positive electrode and the negative electrode; and a non-aqueous electrolyte containing at least one selected from the group consisting of LB(CF₃)₂, LBF₂(CF₃)₃, LBF₂(CF₃), LBC₇(F₃)₄, LBC₇(F₃)₄, LBC₇(F₃)₄, LBC₇(F₃)₄, LBC₇(F₃)₄, LBC₇(F₃)₅, LBC₇(F₃)₅

[0017] The electric double layer capacitor preferably comprises a pair of polarizable electrodes made of activated carbon; as esparatior inference between the electrodes; and a non-aqueous electrody containing at less tone selected from the group consisting of (CH-)₂/NB(CF₂)₂, (CH-)₃/NBF(CF₃)₂, (CH-)₃/NBF(CF₃)₂, (CH-)₃/NBF(CF₃)₂, (CH-)₃/NBF(CF₃)₂, (CH-)₃/NBF(CF₃)₂, (CH-)₃/NBF(CF₃)₂, (CH-)₃/NBF(CF₃)₂, (CH-)₃/NBF(CF₃)₃, (CH-)₃/

[0018] The electroyle capacitor preferably comprises a positive electrode foil having a dielectric layer, a negative electrode foil, as expantor interpresed between the positive electrode foil and the negative electrode foil; and a non-squeous electroyle containing at least one selected from the group consisting of (CH₃)/NBC(G₂)₃, (CH₃)/NBC(

[0019] Generally, a solute is composed of an anion and a cation, and they form an ionic bond. Therefore, the general formula (1): MBR¹R²R³R⁴ may also be expressed as the general formula (1'): M*-(BR¹R²R³R⁴).

[0020] While the novel features of the invention are set forth particularly in the appended claims the invention, both as to organization and content, will be better understood and appreciated, along with other objects and features thereof, from the following detailed description taken in conjunction with the drawings.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWING

[0021] FIG. 1 is a vertical sectional view showing a cylindrical non-aqueous electrolyte battery of the present inven-

[0022] FIG. 2 is a diagram showing a circuit of a device used for evaluating the frequency characteristics of an electrolytic capacitor.

[0023] FIG. 3 is a graph showing the change in the potential difference between the both electrode foils of an electrolytic capacitor of the present invention on which a pulse voltage is applied.

DETAILED DESCRIPTION OF THE INVENTION

[0024] The present invention uses a solute which has a thermal stability substantially equal to that of LIBF, and has an anion portion having a high electronegativity, and easily dissociates into ions in a non-aqueous solvent. Examples of the solute include ones having one or more perfluoroalkyl group represented by the general formula: CnF2n+1 or perfluoroalkylsulfonic acid group represented by the general formula: CmF2m+1SO2 bound to the boron atom. Since a perfluoroalkyl group and a perfluoroalkyl sulfonic acid group have a strong electron withdrawing property, a solute having either of these groups easily dissociates into ions in a non-aqueous solvent. Accordingly, the use of the above solutes can provide a non-aqueous electrolyte having a high thermal stability as well as a high ionic conductivity and, furthermore, it can provide a non-aqueous electrochemical device capable of conducting an efficient high-rate charging and

[0025] It is considered that the more the number of the electron withdrawing groups bound to a boron atom of a solute is, the greater the electronegativity of the anion portion of the solute and the tendency thereof to dissociate into ions become. It is also considered that the more the number of the electron withdrawing groups bound to a boron atom of a solute is, the greater the anionic diameter of the solute becomes and the smaller the tendency thereof to cause an association becomes.

Embodiment 1

[0026] In this embodiment, descriptions will be made on a preferred non-aqueous electrolyte for use in a lithium primary battery, a lithium secondary battery, a lithium ion battery, a polymer battery and the like, A non-aqueous electrolyte of this embodiment can be obtained by dissolving the following solute in the following non-aqueous solvent.

(i) Solute

[0027] A non-aqueous electrolyte of this embodiment contains a solute represented by the general formula; M'BR1R2R3R4, wherein M' is an alkali metal atom such as Li, Na or K and R1 to R4 are electron withdrawing groups or electron withdrawing atoms bound to a boron atom where at least one of R1 to R4 is other than a fluorine atom. [0028] It is preferred that at least one of R1 to R4 is a group represented by the general formula: CnF2nut, where n is an integer of 1 to 4, or a group represented by the general formula: CmF2ma1SO2, where m is an integer of 1 to 4.

The remainders are preferably fluorine atoms.

[0029] Since an anion having a group represented by the general formula: CnF2n+1 is smaller than an anion having a group represented by the general formula: CmF2m+1SO2, a non-aqueous electrolyte containing the former anion has a higher ionic conductivity than the latter one. Therefore, the group represented by the general formula: CoF2001, where n is an integer of 1 to 4 is more preferred than the group represented by the general formula: $C_mF_{2m+1}SO_2$, where m is a integer of 1 to 4.

[0030] The number of the electron withdrawing group other than a fluorine atom may be one or more; however, it is more preferably 2 to 3 from the viewpoint of, for example, ease of the synthesis of the solute. Preferred examples of the solute include solutes represented by the general formula: $M'B(C_nF_{2n+1})_2F_2$ or solutes represented by the general formula: M'B(C_pF₂₀₊₁)₃F.

[0031] Although n and m may be an integer of 1 to 4, it is more preferred that they are 2. The reason is that the electron withdrawing effect of the electron withdrawing group decreases when n and m are too small, and the anionic diameter increases when n and m are too large.

[0032] Specific examples of the solute include LiB(CF₃)_A, LiBF₂(CF₃)_B, LiBF₃(CF₃), LiBF₃(CF₃ $(C_2F_5)_3$, LiBF₂ $(C_2F_5)_2$, LiBF₃ (C_2F_5) , LiB(CF₃SO₂)₄, LiBF(CF₃SO₂)₃, LiBF₂(CF₃SO₂)₂, LiBF₃(CF₃SO₂), LiB $(C_2F_5SO_2)_4$, LiBF $(C_2F_5SO_2)_3$, LiBF $_2(C_2F_5SO_2)_2$, LiBF $_3(C_2F_5SO_2)$ and the like. They may be used singly or as a mixture of two or more of them.

[0033] The non-aqueous electrolyte may further contain LiClO₄, LiBF₆, LiPF₆, LiAlCl₄, LiSbF₆, LiSCN, LiCF₃SO₃, LiCF3CO2, Li(CF3CO2)2, LiAsF6, LiB10Cl10, a lithium lower aliphatic carboxylate, LiCl, LiBr, Lil, chloroboran lithium,

borates such as bis(1,2-benzenediolate(2-)-O,0) lithium borate, bis(2,2-benjethtelenediolate(2-)-O,0) lithium borate, bis(2,2-biphenyidiolate(2-)-O,0) lithium borate and bis(5-fluoro-2-olate-1-benzenesulfonic acid-O,0) lithium borate, imide salts such as bistetrafluoromethanesulfonic acid imide lithium ($(G_7SO_3)AU$), totafluoromethanesulfonic aidi nonafluorobutanesulfonic acid imide lithium ($UN(G_7SO_2)AU$), and bispentafluoroethanesulfonic acid imide lithium ($(G_7SO_3)AU$).

(ii) Non-aqueous solvent

[0034] Examples of the non-aqueous solvent include: cyclic carbonic acid esters such as othylene carbonate, propylene carbonate, butlylene carbonate and winylene carbonate and diversitives thereof; non-cyclic carbonate and sters such as dimethyl carbonate, diethyl carbonate and derly methyl carbonate and derly attention and derivatives thereof; cyclic carbonytic acid esters such as y-butyrolicatione and y-valerolatene and derivatives thereof; alphatic acrbonytic acid esters such as methyl formate, methyl acotate, methyl propionate and ethyl propionate and derivatives thereof; non-cyclic others such as dimethoxyethane, diethyl ether, 1,2-diethoxyethane, ethoxymethoxyethane and 1,3-dioxolane and derivatives thereof; cyclic others such as tetrahydrofuran, 2-methyltetrahydrofuran and 1,3-dioxolane and derivatives thereof; other solvents such as dimethylsulfoxide, formandide, diethylicomandide, dioxolane, acetonitrile, propylintille, nitromethane, ethyl monoglyme, phosphoric acid triesters, trimethcoxymethane, dioxolane derivatives, sulfolane, methylsulfoxide, 1,3-dimethyl-2-midzolidinion,3-methyl-2-oxazolidicanone, ethyl ether, 1,3-propanesultone, anisole and N-methylpyrrolidone; and the like. They may be used singly or as a mixture of two or more of them of the solvential and the solvential according to the solvential a

[0035] Although the concentration of the solute in the non-aqueous electrolyte is not particularly limited, it is preferably 0.2 to 2 mol/liter, and more preferably 0.5 to 1.5 mol/liter.

Embodiment 2

[0036] In this embodiment, descriptions will be made on a non-aqueous electrolyte battery comprising the non-aqueous electrolyte of Embodiment 1. This battery comprises a positive electrode, a negative electrode, a separator interposed between the positive electrode and the negative electrode; and the above non-aqueous electrolyte.

(i) Negative electrode

[0037] The negative electrode is fabricated, for example, by applying onto the surface of a current collector, a negative electrode mixture containing a negative electrode active material, a conductive agent, a binder and the like.

[0038] As the negative electrode active material, for example, lithium metal or a material capable of absorbing and desorbing lithium is used.

[0039] Examples of the material capable of absorbing and desorbing lithium include: the maily decomposed carbon; cokes such as pitch coke, needle coke and portoleum coke; graphites; glassy carbons; phonol resism, materials obtained by sintering and carbonizing furan resins or the like at an appropriate temperature; carbon materials such as carbon fibers and activated carbons; polymer materials such as polyacytene, polypyrrole and polyacene; lithium-containing transition metal oxides such as LLa_M,N, acpailed for the containing transition metal oxides such as LLa_M,N, M, Mg.S and MSit; and tikhum-containing nitrides used as LLa_M,N, where M is a transition metal. They may be used singly or as a mixture of two or more of them. Among them, carbon materials capable of absorbing and desorbing alkali metal to have most commontly used.

[0040] Examples of the conductive agent in the negative electrode include: natural graphites such as flake graphitic, artificial graphities; carbon blacks such as acotylene black, ketchen black, channel black, turnace black, lamp black and thermal black; conductive fibers such as a cropnen fibers and metal fibers; cachon fluoride; metal prowders such as a copper powder and a nickel powder, organic conductive materials such as polyphenylene derivatives. They may be used singly or as a mixture of two or more of them. Among them, artificial graphites, acetylene black and carbon fibers are particularly preferred.

[0041] The amount of the conductive agent used in the negative electrode mixture is preferably 1 to 50 parts by weight, and more preferably 1 to 50 parts by weight of the negative electrode active material. Here, it is not necessary to use the conductive agent when the negative electrode active material has an electronic conductive.

[0042] Examples of the binder in the negative electrode include polyethylene, polypropylene, polyterafluoroethylene, polytriynidene flouride, styrene butadene rubber, totrafluoroethylene-hoxafluorogropylene copolymers, tetrafluoroethyleneylene-perfluorosilly vinyl ether copolymers, vihyildene fluoride-hexafluoropropylene copolymers, vinylidene fluoridechiorotrifluoroethylene copolymers, delylene-tetrafluoroethylene copolymers, polyhortorifluoroethylene, vinylidene

fluoride-pentalfluoropropylene copolymers, propylene-tertafluoroethylene copolymers, ethylene-chiorotrifluoroethylene copolymers, vinylidene fluoride-perfluoromethyl ene copolymers, vinylidene fluoride-perfluoromethyl ene copolymers, ethylene-methyl ene copolymers, ethylene-methyl acrylia copolymers, ethylene-methyl acryliate copolymers and ethylene-methyl methacrylate copolymers. They may be used singly or as a mixture of two or mere of them. Among them, styrene buttadiene rubber, polyvnylidene fluoride, ethylene-acrylic acid copolymers, ethylene-methyl methacrylate copolymers, ethylene-methyl methacrylate copolymers, ethylene-methyl methacrylate copolymers, ethylene-methyl methacrylate copolymers and ethylene-methyl methacrylate copolymers and ethylene-methyl methacrylate copolymers and ethylene-methyl methacrylate copolymers and ethylene-methyl methacrylate copolymers are successful end of experimental ethylene-methylene-methyl methacrylate copolymers are successful end of experimental ethylene-met

[0043] As the current collector in the negative electrode, for example, one made of a stainless steel, nickel, copper, a copper alloy, Italianium, carbon or a conductive polymer is used. Among them, a current collector made of copper or a copper alloy is particularly preferred. The surface of the current collector may be coated with carbon, nickel, Italianium or the like, or if may also be oxidized. It is preferred to form asperties on the surface of the current collector may be in the form of a foil, Iflinn, sheet, net, punched sheet, last sheet, prorous sheet, foamed sheet, nonwown fabric or the like. Generality, the trickness of the current collector in the negative electrods is 1 to 500 mm.

5 (ii) Positive electrode

[0044] The positive electrode is fabricated, for example, by applying onto the surface of a current collector, a positive electrode mixture containing a positive electrode active material, a conductive agent, a binder and the like.

[0045] Examples of the positive electrode active material include $L(COO_3, L(NO_3, L(NO_3, L(CO,N)_1, OO_3, L(NO_3, L(NO_3,$

[0046] Examples of the conductive agent in the positive electrode include: natural graphites such as flake graphite; artificial graphites; carbon blacks such as acetylene black, texten black, tampo black, tampo black, and thermal black; conductive fibers such as carbon fibers and metal fibers; carbon fluoride; metal powders such as a copper, nickel, aluminum or silver provider; conductive misterials such as provide and polisasium titerate; conductive metal oxides such as title such as the such as title such as the such as possible such as the such as t

10047] Examples of the binder in the positive electrode include polyethylene, polypropylene, polyterplationcethylene, polypropylene, polyterplationcethylene expolyments, testraliurocethylene expolymens, testraliurocethylene expolymens, testraliurocethylene expolymens, testraliurocethylene expolymens, testraliurocethylene expolymens, polyteine fluoride-hexafluoropropylene expolymens, vinylidene fluoride chiorottifluorocethylene expolymens, propylene-lettraliurocethylene expolymens, polyteine fluoride-hexafluoropropylene expolymens, propylene-lettraliurocethylene expolymens, vinylidene fluoride-hexafluoropropylene-lettraliurocethylene expolymens, vinylidene fluoride-hexafluoropropylene-actraliurocethylene expolymens, vinylidene fluoride-perfluoromethylene expolym

[0043] As the current collector in the positive electrode, for example, one made of a stainless steel, aluminum, an aluminum alloy, titanium, carbon or a conductive polymer may be used. Among them, a current collector made of aluminum or an aluminum elloy is preferred. The surface of current collector may be coasted with carbon, titanium or the like, or it may also be oxidized. It is preferred to form aspertities on the surface of the current collector. The current collector may be in the form of a foil, film, sheet, net, punched sheet, lath sheet, porous sheet, foamed sheet and nontwors flabric. Generally, the thickness of the current collector in the positive electrode is 1 to 500 miles.

[0049] The positive electrode mixture or the negative electrode mixture may contain various additives, in addition to the above-described materials. They may also contain) a get comprising a polymer material retaining a non-aqueous electrolive.

55 (iii) Separator

[0050] As the separator, an insulating microporous thin film having a high ionic permeability and a predetermined mechanical strength may be used. Additionally, it is preferred to give the separator the capability to increase the re-

sistance by closing its pores in a high temperature range. In terms of the hydrophobic property and the resistance to an organic solvent, it is preferred to use a sheet, a nonwoven tablic or woven fabric made of an olderin type polymer such as polypropylene or polyethylene or a glass fiber. The pore size of the separator is preferably small enough to prevent the positive electrode material, regative electrode material, binder and conductive agent from passing therethrough, each of which is released from the electrode. The pore size may be, for example, 0.01 to 1 jun. Generally, the thickness of the separator is 10 to 300 jun and the porosity thereof is preferably 30 to 80%. Further, a battery can be confligured by integrally modifing a positive electrode and a negative electrode, with a prorus separator made of a polymer material retaining a non-aqueous electrolyte interposed therebetween. As the polymer material, a copolymer of vinyildene flooride and hexaffloropropylene is preferred.

10 [0051] The non-aqueous electrolyte battery of the present invention may be a coin, button, sheet, laminated, cylindrical, flat, or square type.

[0052] The non-aqueous electrolyte battery of the present invention is applicable to, for example, personal digital assistants, portable electronic appliances, domestic power storage devices, two-wheeled motor vehicles, electric vehicles, tythoid electric vehicles, they and the like.

Embodiment 3

16

[0053] In this embodiment, descriptions will be made on a preferred non-aqueous electrolyte for use in an electrolytic capacitor. A non-aqueous electrolyte of this embodiment can be obtained by dissolving the following solute in the following non-aqueous solvent.

(i) Solute

- [0054] A non-equeous electrolyte of this embodiment contains a solute represented by the general formula: M*BR1R2R3R4, wherein M*is an ammonium group and R1 to R4 are electron withdrawing groups or electron withdrawing atoms bound to a boron atom where at least one of R1 to R4 is other than a fluorine atom.
 - [0055] It is preferred that at least one of R¹ to R⁴ is a group represented by the general formula: C_nF_{2n+1}, where n is an integer of 1 to 4, or a group represented by the general formula: C_nF_{2n+1}SO₂, where m is an integer of 1 to 4. The remainders are oreferably fluorine atoms.
- 30 [0056] The group represented by the general formula: C_aF_{2n+1} where n is an integer of 1 to 4, and the group represented by the general formula: C_mF_{2n+1}SO₂, where m is an integer of 1 to 4 are the same as those in Embodiment 1. Therefore, a solute represented by the general formula: M*B(C_mF_{2n+1})₂F₂ and a solute represented by the general formula: M*B(C_mF_{2n+1})₂F₂ are more preferred from the viewpoint of, for example, ease of the synthesis of the solute. (2003T) If it is preferred that the ammonium group M* is represented by the general formula: M*PSPBP7R*B, where R* to
 - The preserve unable are made and independently haloges atoms, alky groups, alkery groups or any groups. Each of R⁸ are bound to N and are each independently haloges atoms, alky groups, alky groups, alky groups, alky groups, alky and the service of R⁸ to R⁸ is preferably an alky group, and more preferably an alky group having 1 to 4 carbon atoms. For example, an ammonium group represented by the general formula: N(C₄H_{26x1})₄, where k is an integer of 1 to 4 is preferred. [0058] Specific examples of the solute include (CH₃),NB(C₃)₄, (CH₃),NB(C₃), (CH₃),NB(C₃)
 - (CF₃), (CH₃),NBF₆(CF₃), (CH₃),CNBF₆(CF₃), (CH₃),CNBF₆(CF₃), (CH₃),CNBF₆(CF₃), (CH₃),CNBF₆(CF₃), (CH₃),CNBF₆(CF₃), (CH₃),CNBF₆(CF₃), (CH₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃), (CH₃),NBF₆(CF₃),NBF₆(CF₃), (CH₃)
 - axhibits a small delectric loss and an excellent charged/discharge characteristic.
 [0059] The non-aqueous electrolyte may further contain MCIO₄, MBF₆, MACI₄, MSF₆, MSCN, MCF₃SO₃, MCF₅CO₂, MKF₆CO₂, MMF₆ MI₁₀O₁₀, MCI, MIR, MI, where M is an ammonium group, quaternary ammonium salts of a lower allphatic carboxylic acid, quaternary ammonium salts of chloroboran, imide saits such as bitetrarfluor-omethanesulfonic acid imide lithium (CF₆SO₂O₂MI). It carfaultorenthanesulfonic acid mide lithium (CF₆SO₂O₂MI), Iterafluoromethanesulfonic acid mide lithium (CF₆SO₂O₂MI). Iterafluoromethanesulfonic acid
- 55 imide lithium (LiN(CF₃SO₂)(C₄F₉SO₂)) and bispentafluoroethanesulfonic acid imide lithium ((C₂F₅SO₂)₂NLi).

(ii) Non-aqueous solvent

[0060] Preferred examples of the non-aqueous solvent include polar protic organic solvents and polar aprotic organic solvents. They may be used singly or as a mixture of two or more of them.

[0061] Examples of the polar profic organic solvent include: monohydric alcohols such as othanol, propend, butanol, organization, cyclopatanol, ocyclopatanol and benzyl alcohol polyhydric alcohols such as othanol, propendio, propindio glycol, grand, cyclopatanol and and benzyl alcohol, polyhydric alcohols such as othanol propindio glycol glycerin, methyl cellosolve, ethyl cellosolve, methoxypropylene glycol and dimethoxypropanol; oxyalcohol compounds; and the like. They may be used simply or as a mixture of two or more of them.

[0062] Examples of the polar aprolic organic solvent include: amide compounds such as N-methylfomamide, N-Modification (N-Modification), N-deptylfomamide, and N,N-dishylfomamide, y-quice activation (acid exters such as 2-butyrioscton) and y-valerolactone; cyclic carboyis class states as ethylene carbonate, propylene carbonate, butylene carbonate and vinylene carbonate; other solvents such as eithylene carbonate, propylene carbonate; other solvents such as eithylene carbonate; other solvents such as either solvents such a

[0063] Although the amount of the solute used is not particularly limited, it is preferably 0.1 to 40 parts by weight per 100 parts by weight of the non-aqueous solvent.

[0064] The voltage proof of the capacitor can be improved by adding bonic acid, a complex compound of boric acid and a polysaccharide (e.g., mannitol or sorbitol), a complex compound of boric acid and a polyhydric alcohol (e.g., ethylene glycof or glycerin), a surfactant, colloids silica or the like in the non-aqueous electrolym.

[0055] Further, an aromatic nitro compound such as p-nitrobenzoic acid or p-nitrophenoi; a piceprorus compound, as uch as p-nitrophenoi acid, pelsperiorus acid, polsperiorus acid, pols

Embodiment 4

[0066] In this embodiment, descriptions will be made on an electrolytic capacitor comprising the non-aqueous electrolyte of Embodiment 3. The electrolytic capacitor comprises a positive electrode (oil having a delectric layer, a negative electrode foil; a separator interposed between the positive electrode foil and the negative electrode foil; and the above non-aqueous electrody.

[0067] The electrolytic capacitor comprises an element obtained by rolling up a positive electrode foil and a negative electrode foil with a separator interposed therebetween; a non-equeous electrolyte with which the capacitor element is impregnated; and a case for accommodating them. The opening of the case is sealed with a sealing member.

[0088] As the positive electrode foil, for example, an aluminum foil is used. A dielectric layer can be formed on the positive electrode foil, for example, by ething the positive electrode foil to expand the surface area finered in a boxic acid and applying a positive potential of 300 to 600 V to the positive electrode foil versus the counter electrode. Through such a step, an oxide lift used as the delectric layer is formed on the positive electrode foil.

[0069] As the negative electrode foil, for example, an aluminum foil is used.

[0070] As the separator, for example, a nonwoven or woven fabric made of a kraft pulp fiber is used.

[0071] Hereinbelow, the present invention will be specifically described with reference to examples, but it should be understood that the invention is not construed as being limited thereto.

EXAMPLE 1

[0072] Each of solutes listed in Table 1 was dissolved at a concentration of 1.0 mol/lifer in a mixed solvent containing ethylence carbonate and ethylence thylence are actionate at a volume ratio of 1.3 thereby to prepare each of electrolytes A to R. Table 1 shows the ionic conductivities at 20 °C of the respective electrolytes measured by using a bridge-type conductivity meter. Here, the ionic conductivities of electrolytes 0 and R containing LIBF₆ and LIPF₆, respectively, at a concentration of 1.0 mol/liter, were also measured for comparison.

| [Table 1] | | | | | |
|---|-------------------------------------|------|--|--|--|
| Electrolyte Solute Ionic conductivity (mS/c | | | | | |
| Α | LiB(CF ₃) ₄ | 7.52 | | | |
| В | LiBF(CF ₃) ₃ | 5.11 | | | |

[Table 1] (continued)

| Electrolyte | Solute | lonic conductivity (mS/cm) |
|-------------|---|----------------------------|
| С | LiBF ₂ (CF ₃) ₂ | 4.02 |
| D | LiBF ₃ (CF ₃) | 3.21 |
| E | LiB(C ₂ F ₅) ₄ | 8.63 |
| F | LiBF(C ₂ F ₅) ₃ | 6.89 |
| G | LiBF ₂ (C ₂ F ₅) ₂ | 4.62 |
| Н | LiBF ₃ (C ₂ F ₅) | 3.31 |
| - 1 | LiB(CF ₃ SO ₂) ₄ | 8.55 |
| J | LiBF(CF ₃ SO ₂) ₃ | 7.62 |
| к | LiBF ₂ (CF ₃ SO ₂) ₂ | 5.83 |
| L | LiBF ₃ (CF ₃ SO ₂) | 4.47 |
| М | LiB(C ₂ F ₅ SO ₂) ₄ | 8.21 |
| N | LiBF(C ₂ F ₅ SO ₂) ₃ | 7.18 |
| 0 | LiBF ₂ (C ₂ F ₅ SO ₂) ₂ | 5.23 |
| P | LiBF ₃ (C ₂ F ₅ SO ₂) | 3.98 |
| Q | LIBF ₄ | 2.91 |
| R | LIPF ₆ | 8.52 |

[0073] From the results shown in Table 1, it can be seen that the use of a solute having one or more perfluoroally/i group or perfluoroally/sulfonic acid group gives a higher ionic conductivity to the non-equeous electrolyte than the use of LIBF₄.

EXAMPLE 2

10

15

20

25

45

[0074] The potential window of each solute was measured by using the electrolytes A to R prepared in Example 1. [0075] Test cells were assembled by using reference electrodes, counter electrodes, test electrodes and the electrolytes A to R. Here, a lithium foil was used for the reference electrode and the counter electrode, and a platinum plate having a surface area of 1 cm² was used for the test electrode.

[0079] Subsequently, the potential of each test electrode was scanned at 10 m/kses to the oxidation direction. The potential of the test electrode versus the potential of the electrode when a current of 100 μ/kcm² was passed was regarded as an oxidation docomposition potential. Meanwhile, the potential of each test electrode was scanned at 10 m/kses to the reduction direction. The potential of the test electrode versus the potential of the reference electrode when a current of 100 μ/kcm² was passed was regarded as a reduction decomposition potential. The results are shown in Table 2.

ITable 21

| | [| |
|---|---|--|
| Solute | Reduction decomposition potential (V) | Oxidation decomposition potential (V) |
| LiB(CF ₃) ₄ | <0.00 | 6.48 |
| LiBF(CF ₃) ₃ | <0.00 | 6.01 |
| LiBF ₂ (CF ₃) ₂ | <0.00 | 5.83 |
| LiBF ₃ (CF ₃) | <0.00 | 5.67 |
| LiB(C ₂ F ₅) ₄ | <0.00 | 6.32 |
| LiBF(C ₂ F ₅) ₃ | <0.00 | 6.11 |
| LiBF ₂ (C ₂ F ₅) ₂ | <0.00 | 5.94 |
| | LIB(CF ₃) ₄ LIBF(CF ₃) ₃ LIBF ₂ (CF ₃) ₂ LIBF ₃ (CF ₃) LIBC ₂ F ₅) ₄ LIBF(C ₂ F ₅) ₃ | $\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$ |

Cable 21 (continued)

| Electrolyte | Solute | Reduction decomposition potential (V) | Oxidation decomposition potential (V |
|-------------|---|---------------------------------------|--------------------------------------|
| н | LiBF ₃ (C ₂ F ₅) | <0.00 | 5.71 |
| 1 | LiB(CF ₃ SO ₂) ₄ | <0.00 | 5.96 |
| J | LiBF(CF ₃ SO ₂) ₃ | <0.00 | 5.89 |
| К | LiBF ₂ (CF ₃ SO ₂) ₂ | <0.00 | 5.66 |
| L | LiBF ₃ (CF ₃ SO ₂) | <0.00 | 5.37 |
| М | LiB(C ₂ F ₅ SO ₂) ₄ | <0.00 | 6.10 |
| N | LiBF(C ₂ F ₅ SO ₂) ₃ | <0.00 | 5.98 |
| 0 | LiBF ₂ (C ₂ F ₅ SO ₂) ₂ | <0.00 | 5.73 |
| Р | LiBF ₃ (C ₂ F ₅ SO ₂) | <0.00 | 5.49 |
| Q | LiBF ₄ | <0.00 | 6.04 |
| R | LiPF ₆ | <0.00 | 6.45 |

[0077] In Table 2, the potential window of each solute having one or more perfluoroally/i group or perfluoroally/isulfonic acid group is substantially equal to or wider than that of LIBF₄. This demonstrates that these solutes are suitable for a non-aqueue electrofyth eather such as a lithium batteru.

EXAMPLE 3

20

25

[0078] Cylindrical lithlum batteries A to R were fabricated by using the electrolytes A to R prepared in Example 1, respectively. FIG. 1 shows a vertical sectional view of the battery thus produced.

[0079] A positive electrode slurry was applied onto a positive electrode current collector made of an aluminum foil, which was then dried and rolled thereby to produce a positive electrode plate 1. The positive electrode slurry was prepared by dispersing in dehydrated N-methylpyprrollidinone, a mixture of 85 parts by weight of a lithium cobattate powder, 10 parts by weight of a carbon powder as a conductive agent and 5 parts by weight of polyvinylidene fluoride as a binder.

[0080] A negative electrode slurry was applied onto a negative electrode current collector made of a copper foil, which was then died and rolled thereby to produce a negative electrode plate 2. The negative electrode slurry was prepared by dispersing in dehydrated N-methylyproficiance, a mixture of 75 parts by weight of a mittical graphic powder, 20 parts by weight of a carbon powder as a conductive agent and 5 parts by weight of polyvinylidene fluoride as a binder.

[0081] The positive electrode plate 1 and the negative electrode plate 2 were rolled up together, with a separator 3 made of polyatily/ene interposed therebetween, thereby to obtain an electrode assembly, husulating rings 8 and 7 were provided on the top and bottom portions of the electrode assembly, respectively. The electrode assembly thus obtained was accommodated in a battery case 8 made of a stainless steel. Then, a positive electrode lead 4 made of a luminum, which was attached to the positive electrode plate 1, was connected to a positive electrode terminal of plate 2, was connected to the bottom of the battery case 8. After a predetermined electrolyte was injected into the battery case 8, the opening of the battery case 8 was sealed with the sealing plate 9 thereby to obtain a battery.

[0082] Each battery thus obtained was 18 mm in diameter and 65 mm in height with a nominal capacity of 1800 mAh. This battery was charged at a contant voltage of 4.2 V, and then discharged at a current of 360 mA or 3600 mA. The discharge capacities thus obtained are shown in Table 3.

[0083] After being discharged at a current of 380 mA, each battery was charged again at a constant voltage of 4.2 V. The charged battery was stored at 80 °C for one month, and was discharged at a current of 3600 mA thereafter. The discharge capacities thus obtained are also shown in Table 3.

(Table 3)

| | | | | Table 3 | | |
|----|---------|-------------|------------------------------------|-------------------------------------|---|--|
| 5 | Battery | Electrolyte | Discharge capacity at 360 mA (mAh) | Discharge capacity at 3600 mA (mAh) | Discharge capacity at 3600 mA after storage (mAh) | |
| 5 | A | Α | 1407 | 1052 | 875 | |
| | В | В | 1318 | 960 | 851 | |
| | С | С | 1297 | 920 | 805 | |
| 10 | D | D | 1223 | 866 | 749 | |
| | E | E | 1638 | 1263 | 1127 | |
| | F | F | 1417 | 1002 | 888 | |
| 15 | G | G | 1345 | 953 | 841 | |
| | н | Н | 1240 | 885 | 766 | |
| | 1 | 1 | 1622 | 1238 | 1217 | |
| 20 | J | J | 1582 | 1194 | 1041 | |
| | к | к | 1354 | 997 | 876 | |
| | ٦ | L | 1305 | 927 | 812 | |
| | М | М | 1518 | 1179 | 1003 | |
| 25 | Ν | N | 1429 | 1058 | 912 | |
| | 0 | 0 | 1332 | 978 | 857 | |
| | Р | Р | 1285 | 904 | 789 | |
| | Q | ď | 1105 | 843 | 432 | |
| 30 | R | R | 1595 | 1210 | 1090 | |

[0084] From the result shown in Table 3, it can be seen that the present invention provides a highly reliable lithium battery which exhibits excellent high-rate discharge characteristics as well as high temperature storage characteristics.

EXAMPLE 4

45

50

55

[0085] Non-aqueous electrolytes 1 to 27 were prepared by dissolving solutes listed in Table 4 at a concentration of 1.0 moi/liter in a mixed solvent containing ethylene carbonate and ethylmethyl carbonate at a volume ratio of 1:3. Table 4 shows the ionic conductivities of the respective electrolytes at 20 °C measured by using the bridge-type conductivity meter. Here, the ionic conductivities of electrolytes 25 to 27 containing (CH₃)₄NBF₄, (C₂H₅)₄NBF₄ and ((CH₃)₃C)₄NBF₄, respectively, at a concentration of 1.0 mol/liter, were also measured for comparison.

Table 41

| [lable 4] | | | | |
|-------------|--|----------------------------|--|--|
| Electrolyte | Solute | lonic conductivity (mS/cm) | | |
| 1 | (CH ₃) ₄ NB(CF ₃) ₄ | 7.22 | | |
| 2 | (CH ₃) ₄ NBF(CF ₃) ₃ | 5.02 | | |
| 3 | (CH ₃) ₄ NBF ₂ (CF ₃) ₂ | 3.98 | | |
| 4 | (CH ₃) ₄ NBF ₃ (CF ₃) | 3.09 | | |
| 5 | (CH ₃) ₄ NB(C ₂ F ₅) ₄ | 8.31 | | |
| 6 | (CH ₃) ₄ NBF(C ₂ F ₅) ₃ | 6.72 | | |
| 7 | (CH ₃) ₄ NBF ₂ (C ₂ F ₅) ₂ | 4.43 | | |
| 8 | (CH ₃) ₄ NBF ₃ (C ₂ F ₅) | 3.27 | | |
| 9 | (C ₂ H ₅) ₄ NB(CF ₃) ₄ | 6.88 | | |

EP 1 174 941 A2

(Table 4) (continued)

| Electrolyte | Solute | Ionic conductivity (mS/cm) | |
|-------------|---|----------------------------|--|
| 10 | (C ₂ H ₅) ₄ NBF(CF ₃) ₃ | 4.71 | |
| 11 | (C ₂ H ₅) ₄ NBF ₂ (CF ₃) ₂ | 3.60 | |
| 12 | (C ₂ H ₅) ₄ NBF ₃ (CF ₃) | 2.98 | |
| 13 | (C ₂ H ₅) ₄ NB(C ₂ F ₅) ₄ | 7.19 | |
| 14 | (C ₂ H ₅) ₄ NBF(C ₂ F ₅) ₃ | 4.97 | |
| 15 | (C ₂ H ₅) ₄ NBF ₂ (C ₂ F ₅) ₂ | 3.89 | |
| 16 | (C ₂ H ₅) ₄ NBF ₃ (C ₂ F ₅) | 3.01 | |
| 17 | ((CH ₃) ₃ C) ₄ NB(CF ₃) ₄ | 6.19 | |
| 18 | ((CH ₃) ₃ C) ₄ NBF(CF ₃) ₃ | 4.45 | |
| 19 | ((CH ₃) ₃ C) ₄ NBF ₂ (CF ₃) ₂ | 3.32 2.69 | |
| 20 | ((CH ₃) ₃ C) ₄ NBF ₃ (CF ₃) | | |
| 21 | ((CH ₃) ₃ C) ₄ NB(C ₂ F ₅) ₄ | 6.74 | |
| 22 | ((CH ₃) ₃ C) ₄ NBF(C ₂ F ₅) ₃ | 4.38 | |
| 23 | ((CH ₃) ₃ C) ₄ NBF ₂ (C ₂ F ₅) ₂ | 3.26 | |
| 24 | ((CH ₃) ₃ C) ₄ NBF ₃ (C ₂ F ₅) 2.72 | | |
| 25 | (CH ₃) ₄ NBF ₄ | 2.79 | |
| 26 | (C ₂ H ₅) ₄ NBF ₄ | 2.36 | |
| 27 | ((CH ₃) ₃ C) ₄ NBF ₄ | 2.05 | |

[0086] From the results shown in Table 4, it can be seen that the use of each solute having one or more perfluoroalityl group glives a higher ionic conductivity to the non-aqueous electrolyte than the use of the solute whose anion portion is BF4.

35 EXAMPLE 5

10

20

25

40

[0087] The potential window of each solute was measured in the same manner as in Example 2, except for using the non-aqueous electrolytes 1 to 27 prepared in Example 4. The results are shown in Table 5.

[Table 5]

| | Electrolyte | Solute | Reduction decomposition potential (V) | Oxidation decomposition potential (V) |
|----|-------------|--|--|---------------------------------------|
| | 1 | (CH ₃) ₄ NB(CF ₃) ₄ | <0.00 | 6.47 |
| 45 | 2 | (CH ₃) ₄ NBF(CF ₃) ₃ | <0.00 | 6.16 |
| | 3 | (CH ₃) ₄ NBF ₂ (CF ₃) ₂ | <0.00 | 6.04 |
| 50 | 4 | (CH ₃) ₄ NBF ₃ (CF ₃) | <0.00 | 5.95 |
| | 5 | (CH ₃) ₄ NB(C ₂ F ₅) ₄ | <0.00 | 6.34 |
| | 6 | (CH ₃) ₄ NBF(C ₂ F ₅) ₃ | <0.00 | 6.25 |
| | 7 | (CH ₃) ₄ NBF ₂ (C ₂ F ₅) ₂ | <0.00 | 6.11 |
| 55 | 8 | (CH ₃) ₄ NBF ₃ (C ₂ F ₅) | <0.00 | 5.99 |
| | 9 | (C2H ₅) ₄ NB(CF ₃) ₄ | <0.00 | 6.26 |
| | 10 | (C ₂ H ₅) ₄ NBF(CF ₃) ₃ | <0.00 | 6.08 |

Table 51 (continued)

| Electrolyte | Solute | Reduction decomposition potential (V) | Oxidation decomposition potentia (V) |
|-------------|---|---------------------------------------|---|
| 11 | (C ₂ H ₅) ₄ NBF ₂ (CF ₃) ₂ | <0.00 | 5.94 |
| 12 | (C ₂ H ₅) ₄ NBF ₃ (CF ₃) | <0.00 | 5.85 |
| 13 | (C ₂ H ₅) ₄ NB(C ₂ F ₅) ₄ | <0.00 | 6.18 |
| 14 | (C ₂ H ₅) ₄ NBF(C ₂ F ₅) ₃ | <0.00 | 6.13 |
| 15 | (C ₂ H ₅) ₄ NBF ₂ (C ₂ F ₅) ₂ | <0.00 | 5.99 |
| 16 | (C ₂ H ₅) ₄ NBF ₃ (C ₂ F ₅) | <0.00 | 5.90 |
| 17 | ((CH ₃) ₃ C) ₄ NB(CF ₃) ₄ | <0.00 | 6.20 |
| 18 | ((CH ₃) ₃ C) ₄ NBF(CF ₃) ₃ | <0.00 | 5.99 |
| 19 | ((CH ₃) ₃ C) ₄ NBF ₂ (CF ₃) ₂ | <0.00 | 5.86 |
| 20 | ((CH ₃) ₃ C) ₄ NBF ₃ (CF ₃) | <0.00 | 5.77 |
| 21 | ((CH ₃) ₃ C) ₄ NB(C ₂ F ₅) ₄ | <0.00 | 6.11 |
| 22 | ((CH ₃) ₃ C) ₄ NBF(C ₂ F ₅) ₃ | <0.00 | 6.05 |
| 23 | ((CH ₃) ₃ C) ₄ NBF ₂ (C ₂ F ₅) ₂ | <0.00 | 5.92 |
| 24 | ((CH ₃) ₃ C) ₄ NBF ₃ (C ₂ F ₅) | <0.00 | 5.81 |
| 25 | (CH ₃) ₄ NBF ₄ | <0.00 | 6.02 |
| 26 | (C ₂ H ₅) ₄ NBF ₄ | <0.00 | 5.93 |
| 27 | ((CH ₃) ₃ C) ₄ NBF ₄ | <0.00 | 5.82 |

[0088] In Table 5, the potential window of each solute having one or more perfluoroality! group is substantially equal to or wider than that of each solute whose anion portion is BF₄. This demonstrates that these solutes are suitable for electrolytic capacitors.

EXAMPLE 6

10

18

20

25

[0089] Electrolytic capacitors comprising the electrolytes 1 to 27 prepared in Example 4 were produced in the following manner.

[0090] The surface of an aluminum foil having a thickness of 100 µm was etched by electrolysis. A positive potential of 500 V versus the potential of the counter electrode was applied on the etched aluminum foil in a boric acid, which was then stood still for 15 minutes. As a result, an aluminum oxide film used as a dielectric layer was formed on the surface of the aluminum foil.

[0091] Subsequently, an aluminum foil having an aluminum oxide film used as a positive electrode foil and another aluminum foil used as a negative electrode foil were folled up together, with a separator made of a kraft pulp fiber interposed therebetween, thereby to obtain a capacitor element. The capacitor element thus obtained was impregnated with a predetermined non-aqueous electrolyte. The capacitor element impregnated with the non-aqueous electrolyte. The capacitor element impregnated with the non-aqueous electrolyte was accommodated in a case made of aluminum. Then, the opening of the case was seeled with an insultating sealing member, and the leads of the positive electrode foil and the negative electrode foil were threaded through the sealing member to be publied outside, thresho obtaining an electrolytic capacitor.

[0092] Subsequently, the electrolytic capacitor thus obtained was evaluated for the frequency characteristics.

[0083] A circuit as shown in FIG. 2 was assembled, which comprised a direct-current power source 11, an electrolytic capacitor 12, a resister 13, a relate which 14 and a voltment e15. The negative side of the direct-current power source, the negative electrode foil side of the electrolytic capacitor and the resistor were grounded. Then, the relay switch was activated at a duty ratio of 30 (60%) and at a frequency of 60Pt to apply a pulse votage on the electrolytic capacitor. Here, the upper limit of the applied voltage was set at 3.0 V. FIG. 3 shows the change in the difference between the potentials of the negative electrods foil and the possitive electrode foil off the electrolytic capacitor.

[0094] The voltage value of the electrolytic capacitor was recorded 1/240 second after the start of the charging during each charging interval, and this measurement was conducted for 10 seconds. The average values of the recorded

voltages are shown in Table 6. It can be said that the charge/discharge characteristics improve with an increase in the speed of the potential rise on the positive electrode foil.

[Table 6]

| | | [Table 6] | |
|-----------|-------------|---|-------------|
| Capacitor | Electrolyte | Solute | Voltage (V) |
| 1 | 1 | (CH ₃) ₄ NB(CF ₃) ₄ | 2.94 |
| 2 | 2 | (CH ₃) ₄ NBF(CF ₃) ₃ | 2.80 |
| 3 | 3 | (CH ₃) ₄ NBF ₂ (CF ₃) ₂ | 2.59 |
| 4 | 4 | (CH ₃) ₄ NBF ₃ (CF ₃) | 2.28 |
| 5 | 5 | (CH ₃) ₄ NB(C ₂ F ₅) ₄ | 2.96 |
| 6 | 6 | (CH ₃) ₄ NBF(C ₂ F ₅) ₃ | 2.89 |
| 7 | 7 | (CH ₃) ₄ NBF ₂ (C ₂ F ₅) ₂ | 2.66 |
| 8 | 8 | (CH ₃) ₄ NBF ₃ (C ₂ F ₅) | 2.35 |
| 9 | 9 | (C ₂ H ₅) ₄ NB(CF ₃) ₄ | 2.51 |
| 10 | 10 | (C ₂ H ₅) ₄ NBF(CF ₃) ₃ | 2.74 |
| 11 | 11 | (C ₂ H ₅) ₄ NBF ₂ (CF ₃) ₂ | 2.44 |
| 12 | 12 | (C ₂ H ₅) ₄ NBF ₃ (CF ₃) | 2.20 |
| 13 | 13 | (C ₂ H ₅) ₄ NB(C ₂ F ₅) ₄ | 2.93 |
| 14 | 14 | (C ₂ H ₅) ₄ NBF(C ₂ F ₅) ₃ | 2.78 |
| 15 | 15 | (C ₂ H ₅) ₄ NBF ₂ (C ₂ F ₅) ₂ | 2.53 |
| 16 | 16 | (C ₂ H ₅) ₄ NBF ₃ (C ₂ F ₅) | 2.22 |
| 17 | 17 | ((CH ₃) ₃ C) ₄ NB(CF ₃) ₄ | 2.88 |
| 18 | 18 | ((CH ₃) ₃ C) ₄ NBF(CF ₃) ₃ | 2.69 |
| 19 | 19 | ((CH ₃) ₃ C) ₄ NBF ₂ (CF ₃) ₂ | 2.39 |
| 20 | 20 | ((CH ₃) ₃ C) ₄ NBF ₃ (CF ₃) | 2.15 |
| 21 | 21 | ((CH ₃) ₃ C) ₄ NB(C ₂ F ₅) ₄ | 2.90 |
| 22 | 22 | ((CH ₃) ₃ C) ₄ NBF(C ₂ F ₅) ₃ | 2.63 |
| 23 | 23 | ((CH ₃) ₃ C) ₄ NBF ₂ (C ₂ F ₅) ₂ | 2.33 |
| 24 | 24 | ((CH ₃) ₃ C) ₄ NBF ₃ (C ₂ F ₅) | 2.16 |
| 25 | 25 | (CH ₃) ₄ NBF ₄ | 2.18 |
| 26 | 26 | (C ₂ H ₅) ₄ NBF ₄ | 2.11 |
| 27 | 27 | ((CH ₃) ₃ C) ₄ NBF ₄ | 2.02 |
| 27 | 27 | ((CH ₃) ₃ C) ₄ NBF ₄ | 2.02 |

[0095] From the results shown in Table 6, it can be seen that the electrolytic capacitor using each solute having one or more perfluoreality group exhibits superior frequency characteristics. This is presumably due to the increase in the ionic conductify of the non-aqueous electrolytic.

[0096] When a similar measurement was conducted at an increased charge voltage, the electrolytic capacitor using each solute having one or more perfluorability group showed filter detendration in the charge/discharge characteristics. The deterioration of the charge/discharge characteristics was presumably due to the decomposition of an anion by notifiation.

[0097] From the results, it can be seen that the use of each solute having one or more perfluoroalkyl group can improve the stability of an electrolytic capacitor.

[0088] As described above, the present invention uses a solute which has a thermal stability substantially equal to that of LIBF, and an anion portion having a high electronegality, and easily dissociates into ions. Therefore, it provides a non-equeous electrolyte having a high thermal stability as well as a high into conductivity. Furthermore, the use

10

20

48

the non-aqueous electrolyte of the present invention can provide a non-aqueous electrochemical device having an excellent high-rate charge/discharge characteristic.

[0099] Although the present invention has been described in terms of the presently preferred embodiments, it is to be understood that such disclosure is not to be interpreted as limiting. Various alterations and modifications will no doubt become apparent to those skilled in the art to which the present invention pertains, after having read the above disclosure. Accordingly, it is intended that the appended claims be interpreted as covering all alterations and modifications as fall within the true solit and scope of the invention.

[0100] A non-aqueous electrolyte is disclosed, which comprises a non-aqueous solvent and a solute represented by the general formula(1). MBRPRPR, wherein M is an alkall metal atom or a momnnium group and N to R ** are each independently electron withdrawing groups or electron withdrawing atoms bound to B where at least one of R to R ** is other than a fluorine store. The solute has a themat stability substantially equal to that of LIBF, and an anion portion having a high electronegativity, and easily dissociates into ions. Therefore, a non-aqueous electrolyte containing this solute has a high ionic conductivity and is difficult to cause a generation of a gas or deterioration in characteristics due to the decomposition of the solute, which occurs during use at high temperatures or after storage at high temperatures.

Claims

30

35

50

1. A non-aqueous electrolyte, comprising a non-aqueous solvent and a solute represented by the general formula (1):

- 25 wherein M is an alkali metal atom or an ammonium group and R¹ to R⁴ are each independently electron withdrawing groups or electron withdrawing atoms bound to B where at least one of R¹ to R⁴ is other than a fluorine atom.
 - The non-aqueous electrolyte in accordance with claim 1, wherein at least one of R¹ to R⁴ is represented by the general formula (2): C_mF_{2n+1}, where n is an integer of 1 to 4, or the general formula (3): C_mF_{2m+1}SO₂, where m is an integer of 1 to 4.
 - The non-aqueous electrolyte in accordance with claim 1, wherein said solute is at least one selected from the group consisting of LB(ErG₂), LBF(ErG₂), LBF(Er
 - 4. The non-aqueous electrolyte in accordance with claim 1, wherein said ammonium group is represented by the general formula (4): NB²R²R²R², where R² to R² are bound to N and are each independently hydrogen atoms, allyl groups, alkenyl groups or anyl groups.
 - 5. The non-aqueous electrolyte in accordance with claim 1, wherein said solute is at least one selected from the group consisting of (CH₃)/NBE(CF₃)₂, (CH₃)/NBF(CF₃)₂, (CH₃)/NBF(CF₃)₃, (CH₃)/NBF(CF₃)₃,
 - 6. An electrochemical device, comprising the non-aqueous electrolyte in accordance with claim 1.
 - A non-aqueous electrolyte băttery, comprising a positive electrode; a negative electrode; a separator interposed between said positive electrode and said negative electrode; and the non-aqueous electrolyte in accordance with claim 3.

- An electrolytic capacitor, comprising a positive electrode foil having a dielectric layer; a negative electrode foil; a separator interposed between said positive electrode foil and said negative electrode foil; and the non-aqueous electrolyte in accordance with claim 5.
- An electric double layer capacitor, comprising a pair of polarizable electrodes made of activated carbon; a separator
 interposed between said electrodes; and the non-aqueous electrolyte in accordance with claim 5.

10

15

20

30

50

FIG. 1

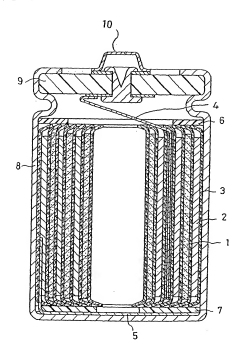
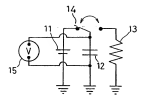


FIG. 2



F I G. 3

INSDOCID: <EP_____1174941A2_L>

